Posterior interosseous neuropathy
Supinator syndrome vs fascicular radial neuropathy

ABSTRACT

Objective: To investigate the spatial pattern of lesion dispersion in posterior interosseous neuropathy syndrome (PINS) by high-resolution magnetic resonance neurography.

Methods: This prospective study was approved by the local ethics committee and written informed consent was obtained from all patients. In 19 patients with PINS and 20 healthy controls, a standardized magnetic resonance neurography protocol at 3-tesla was performed with coverage of the upper arm and elbow (T2-weighted fat-saturated: echo time/repetition time 52/7,020 milliseconds, in-plane resolution 0.27 x 0.27 mm²). Lesion classification of the radial nerve trunk and its deep branch (which becomes the posterior interosseous nerve) was performed by visual rating and additional quantitative analysis of normalized T2 signal of radial nerve voxels.

Results: Of 19 patients with PINS, only 3 (16%) had a focal neuropathy at the entry of the radial nerve deep branch into the supinator muscle at elbow/forearm level. The other 16 (84%) had proximal radial nerve lesions at the upper arm level with a predominant lesion focus 8.3 ± 4.6 cm proximal to the humeroradial joint. Most of these lesions (75%) followed a specific somatotopic pattern, involving only those fascicles that would form the posterior interosseous nerve more distally.

Conclusions: PINS is not necessarily caused by focal compression at the supinator muscle but is instead frequently a consequence of partial fascicular lesions of the radial nerve trunk at the upper arm level. Neuroimaging should be considered as a complementary diagnostic method in PINS. Neurology® 2016;87:1–8

GLOSSARY

CSA = cross-sectional area; MRN = magnetic resonance neurography; PIN = posterior interosseous nerve; PINS = posterior interosseous neuropathy syndrome; RN = radial nerve; ROI = region of interest.
Sonography in supinator syndrome has been reported to detect focal PIN swelling at entry into the supinator muscle.\textsuperscript{10–14} MRN detects nerve lesions as increased T2-weighted signal and helps to discriminate neuromorphologies of compressive origin from other etiologies by the pattern of lesion localization.\textsuperscript{15,16} In this prospective investigation, we sought to determine lesion sites and the spatial pattern of lesion dispersion by MRN in patients with a clinical diagnosis of PINS.

METHODS

Standard protocol approvals, registrations, and patient consents. The study was approved by the institutional ethics committee (S-398/2012). All participants gave written informed consent.

Clinical and demographic patient data. Patients included in the study were referred to our department with a clinical diagnosis of PINS due to muscle weakness and abnormal EMG results in the distribution of the PIN (i.e., extensor digitorum, extensor digiti minimi, extensor carpi ulnaris, abductor pollicis longus, extensor pollicis brevis, extensor pollicis longus, extensor indicis).

Patients with motor weakness involving more proximal muscles (triceps brachii muscle, brachioradialis, extensor carpi radialis longus and brevis) or EMG evidence of denervation in these were excluded from the study. Patients with sensory deficits, especially in the distribution of the superficial sensory RN, were also excluded from the study. Furthermore, patients with nerve root compression (C6-C8) on cervical spine imaging were excluded from the study.

Overall, 28 patients (13 women, 15 men, mean age 48.5 ± 15.5 years) fulfilled the inclusion criteria and were examined at the Department of Neuroradiology of Heidelberg University Hospital, Germany, between January 2010 and December 2015. Of these 28 patients, 9 were excluded after careful review of their history and after electrophysiologic reports revealed findings indicative of a neuropathy other than a compression neuropathy at the level of the supinator muscle (4 with additional sensory RN symptoms, 4 with more proximal muscle paresis or EMG evidence of denervation, one with cervical root compression at C7).

Twenty age- and sex-matched participants (8 women, 12 men, 49.9 ± 16.7 years) without symptoms or signs of radial neuropathy, or risk factors for polyneuropathy such as diabetes, chemotherapy, or infectious diseases, served as control group.

MRN imaging. Examinations were conducted using a 3-tesla unit (MAGNETOM Verio; Siemens AG, Erlangen, Germany) as described previously.\textsuperscript{26} Participants were examined in prone position in a knee 8-channel, phased-array receive coil with the arm in elbow extension. The magic angle effect was avoided by aligning the longitudinal axis of the upper arm at an angle of ±10° relative to the B0 field. Two to 3 image slabs were acquired for large imaging coverage of the RN, its branches, and the muscles it innervates in the forearm: (1) proximal: at the upper arm level; (2) central: at the elbow centered on the supinator muscle; and (in case forearm muscles were not sufficiently covered by sequence 2) by (3) at the forearm.

Sequence measures for the transversal T2-weighted turbo spin echo were as described previously:\textsuperscript{26} repetition time/echo time 6,980/52 milliseconds, spectral fat saturation, slice thickness 3.0 mm, number of slices 45, interslice gap 0.3 mm, field of view 130 × 130 mm², acquisition matrix 512 × 358, pixel spacing 0.254 × 0.254 mm², number of excitations = 3, acquisition time 7:17 minutes.

As an option, an additional sagittal-oblique T2-weighted sequence using a dedicated surface coil (NORAS, Würzburg, Germany) was acquired for assessment of the brachial plexus with the following measures: repetition time/echo time 5,530/45 milliseconds, spectral fat saturation, slice thickness 3.0 mm, number of slices 51, interslice gap 0.3 mm, field of view 150 × 150 mm², acquisition matrix 320 × 198, pixel spacing 0.47 × 0.47 mm², number of excitations = 3, acquisition time 7:46 minutes.

Qualitative image analysis. Images were assessed by 2 neuroradiologists (P.B., M.P.) with more than 7 and 10 years of experience in MRN, respectively, similar to a previous analysis\textsuperscript{17} regarding the following items:

1. Lesion determination: dichotomous consensus ratings regarding the presence or absence of lesions, based on T2-weighted signal increase and caliber of RN and PIN fascicles.
2. Predominant lesion localization: the anatomical position of predominant lesion focus of the PIN or RN, defined as strongest increase in T2-weighted signal and caliber, measured as distance from the humeroradial joint.
3. Fascicular involvement: dichotomous consensus ratings on lesion involvement of the entire nerve cross-section or only a partial cross-sectional area (CSA) (fascicular lesion).
4. Longitudinal involvement: those patients in whom an additional sequence of the brachial plexus was acquired were assessed for the presence of lesions within the brachial plexus cords.

Quantitative image and statistical analysis. Further steps on quantitative analysis were performed on a Siemens Syngo Workstation (SyngoMMWP VE31A, syngVE32B) by 2 investigators (P.B., A.X.) blinded to all participant data. Spatial registration for each participant was performed with reference to the humeroradial joint space. RN T2-weighted contrast was obtained at the site of predominant lesion focus by manual segmentation of the nerve circumference for measurement of signal intensities within this region of interest (nROI) as well as the CSA. For healthy controls, 2 positions were chosen: one 8.3 cm proximal to the humeroradial joint was chosen for this, corresponding to the mean distance of predominant proximal lesion focus in patients with proximal lesions; and another, 1.3 cm distal to the humeroradial joint, corresponding to the nerve entry to the supinator muscle. Additional ROIs were placed on the same slice within the long head of the biceps brachii muscle (mROI).

For each participant, normalized RN T2 values were calculated as follows: nT2 = nROI/mROI. Patient and control values were tested against each other using the 2-sided Student t test, with the significance level defined at $p < 0.05$.

RESULTS

Clinical findings and patient data. Nineteen patients (9 women, 10 men, mean age 48.5 ± 14.7 years) were included in the study with a diagnosis of PINS based on clinical examination and electrophysiologic test results (table).

Eight patients reported having initially experienced transient pain in the upper extremity, without evidence of sensory findings on physical examinations. Six patients had previously undergone operation for a presumed PIN compression syndrome at...
the elbow but had not shown improvement in muscle strength. Sonography had been performed in 10 patients before MRN. In 4, sonographic findings with swelling at the entry into the supinator muscle appeared consistent with a compression neuropathy. In 2, unclear swelling of the RN was found at the upper arm level; in another one, nerve caliber was normal but slight hypoechogenicity of some fascicles was noted at the upper arm level. Three had normal ultrasound findings.

**MRN imaging findings.** None of the healthy controls displayed an increased T2-weighted signal in the RN. In contrast, all of the 19 patients with PINS exhibited abnormal findings on MRN: 3 (15.8%) were found to have a focal neuropathy at the entry to or within the supinator muscle. Patients 2 and 17 were found to have increased signal at the exact point of entry into the supinator muscle. Patient 4 had severe swelling of the nerve within the supinator muscle. Unexpectedly, the other 16 patients (84.2%) showed a distinct pattern with proximally located lesions within the RN (figure 1). All of these were apparent over a larger longitudinal segment of the RN trunk at the upper arm level.

**Lesion localization.** The distance of the predominant lesion focus within the PIN or RN was determined in relation to the elbow. For the 16 patients with proximally extending lesions, this was 8.3 ± 4.6 cm proximal to the humeroradial joint, while for the 3 patients with focal nerve lesions within the supinator muscle, this was 1.3 ± 1.8 cm distal to the humeroradial joint (equivalent to entry to or within the supinator muscle).

Of the 16 patients with a proximal lesion focus, pathologically increased T2-weighted signal was observed in the entire RN cross-section in 4 (25.0%) of them, and no normal-appearing fascicles could reliably be distinguished at the level of muscle. Unexpectedly, the other 16 patients (84.2%) showed a distinct pattern with proximally located lesions within the RN (figure 1). All of these were apparent over a larger longitudinal segment of the RN trunk at the upper arm level.

**Muscle strength is given according to the grading (0–5) by the Medical Research Council. By definition of the inclusion criteria, muscle strength was 5 for all muscle groups not shown in the table.**

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<th>Finger extension</th>
<th>Thumb extension</th>
<th>Wrist extension</th>
<th>Initial pain</th>
<th>Final diagnosis</th>
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predominant nerve lesion focus. The other 12 of 16 patients (75.0%) had partial, fascicular lesions (figure 2). The precise location for these was in all cases at the dorsomedial portion of the RN cross-section, even when lesions extended proximally to the humeral groove. This lesion area coincided with the topographic internal arrangement of motor fascicles within the RN, as originally depicted in a somatotopic map by Sunderland (figure 3). These hyperintense fascicles then continued distally to form the deep branch of the RN, or PIN. In contrast, the anterolateral fascicles were not involved and continued distally to form the superficial branch of the RN.

Lesion extension into the brachial plexus. In 8 of the 16 patients with proximal lesions, the longitudinal lesion extension at the upper arm level led to the acquisition of additional, optional imaging sequences to cover the brachial plexus in the same imaging session. Brachial plexus lesions were detected in 7 patients (87.5%), all of them restricted to the posterior cord from which the RN arises. At the same time, no lesions were detected in the medial or lateral cord (figure 4).

Quantitative analysis. At the site of predominant lesion focus, mean normalized RN T2 signal was increased in patients with proximal lesions compared to controls (11.4 ± 6.2 vs 6.7 ± 1.3 mm²; p < 0.001). The 3 patients with PIN lesions in the supinator muscle had increased CSA (10.7 ± 12.9 vs 3.6 ± 1.4 mm²; p = 0.02) and T2 SR (1.36 ± 0.07 vs 1.11 ± 0.24; p = 0.03) compared to controls at their site of predominant lesion focus in the supinator muscle.

DISCUSSION Patients with isolated finger drop are often primarily assumed to have a focal entrapment neuropathy caused by compression of the deep motor branch of the RN at the supinator tunnel. In this study, we present evidence by neuroimaging that initially unsuspected proximal lesions of the RN trunk at the upper arm level are frequently the cause of this clinical presentation.

The majority of our patients had proximal, longitudinally extending lesions along the upper arm, often selectively involving only dorsomedially located motor fascicles within the RN. In the literature, a number of scattered case reports exist that describe patients with a clinical picture initially consistent with PINS, who upon surgical exploration, had lesions in the RN or PIN at mostly proximal sites without detectable external compression.18–21 A single case report with confirmation by MRN findings was
recently added to these few individual descriptions. Our study is the first to systematically investigate this observation in a larger group of patients with PINS.

How can we account for this apparent discrepancy between clinical reasoning and neuroimaging results? There are 2 potential explanations.

The first explanation is based on the concept of peripheral nerve somatotopy. Peripheral nerves are now known to harbor a high degree of consistent longitudinal arrangement of their fascicles with highly relevant clinical implications for lesion localization in the PNS. If a given proximal nerve lesion involves specifically only those fascicles destined to form a particular branch that leaves the nerve more distally, but does not involve fascicles to other branches, the clinical manifestation of this lesion will mimic a more distal lesion of the particular branch. This is obvious in the case of traumatic partial nerve injuries. In spontaneously occurring neuropathies, it represents a major pitfall in diagnosis as has recently been shown for the anterior interosseous nerve syndrome. Historical dissections of the RN and its branches suggest that the RN is also potentially susceptible to such erroneous localization of a lesion. The observed somatotopic lesion pattern in our patients, in close correspondence to the somatotopic mappings obtained from Sunderland ex vivo, seems to confirm this hypothesis.

The other potential explanation would be a length-dependence of the underlying disease. A length-dependent occurrence of symptoms has been investigated for many polyneuropathies including diabetic polyneuropathy, immunoglobulin M gammopathy–associated polyneuropathy, or multifocal motor neuropathy. Recent neuroimaging studies of diabetic and neurofibromatosis-associated polyneuropathies have shown the disseminated multifocality of nerve lesions whose cumulation correlates with symptom severity. A length-dependent radial mononeuropathy could first present with finger drop, as seen in our patients, before eventually proceeding to involve additional muscles. This was actually the case in 4 patients excluded from our study because their disease had progressed before the MRI to also involve more proximal muscles. Length-dependence therefore seems to be an additional possible explanation.

The longitudinal and disseminated involvement of the RN suggests an inflammatory process as underlying disease entity. Clinical follow-up was not part
of the study design, but several patients were subsequently diagnosed with an inflammatory nerve condition such as multifocal motor neuropathy. The documented extension of lesions into the posterior cord of the brachial plexus in several patients also points to an overlap with brachial plexus neuritis. In fact, certain patients with Parsonage-Turner syndrome are known to present with symptoms pointing toward involvement along the peripheral nerve trunks distal to the level of the brachial plexus. Furthermore, the transient pain at symptom onset that several patients reported is a typical infrequent feature of inflammatory nerve conditions such as multifocal motor neuropathy or neuralgic amyotrophy.35,36

The clinical implication of our study, then, is that neuroimaging including the proximal course of the RN should be considered for patients presenting with finger drop since clinical reasoning alone and EMG cannot always reliably differentiate between sole compression at the elbow and selective fascicular involvement of proximal nerve portions at the upper arm level. This is especially relevant for patients who would otherwise undergo surgical exploration for presumed compression of the nerve at the elbow. Imaging will either allow for more precise planning of the surgical strategy if focal, compressive structures can be identified, or it may render surgery altogether inappropriate. A second interesting implication of the study is the clinical relevance of peripheral nerve somatotopy, which can be delineated by MRN7,17,25 and which was in good agreement with historical dissection findings by Sunderland.29

Ultrasound was not part of the standard protocol in our study. In 3 of the 10 patients who underwent ultrasound examination before MRI, proximal RN abnormalities were also documented on nerve ultrasound. Nerve caliber increase at proximal arm positions can well be detected by ultrasound experts and has been described in the literature to detect RN lesions.37–39 Caliber-neutral lesions are much harder to identify, and MRN likely has better sensitivity for these lesions.37 Generally, ultrasound seems feasible for detection of distal nerve lesions while MRN is likely necessary for far proximal lesions.39

There are limitations to our study. Our patient population should not be considered fully representative for all patients presenting with finger drop since 6 patients had already been operated without

Figure 3 Somatotopy of posterior interosseous nerve fascicular lesions on individual level and group level compared to atlas

(A) The T2-weighted source image of the radial nerve of patient 7 is shown for the site of predominant lesion focus (9.6 cm proximal to the humeroradial joint). Anatomical orientation is given by labeling ventral/dorsal/medial/lateral contours. (B) Spatial map of the patient group mean normalized T2 signal. This cross-sectional lesion area is at the dorsal/posterior and medial aspect of the radial nerve at the upper arm level with a mean distance of 8.3 cm proximal to the humeroradial joint space. The map was rendered after segmentation and intersubject image registration. (C) Somatotopic/topographic internal map of fascicles of the radial nerve trunk, obtained ex vivo as schematic drawing (modified from Sunderland S. The intraneural topography of the radial, median and ulnar nerves. Brain 1945;68:243–299,29 by permission of Oxford University Press). On this map, the fascicles identified as posterior interosseous fascicles (black filled dots with red background) are in spatial arrangement with the T2 lesion focus on individual (A) and group level (B). Note that lesion focus appears larger on the averaged map compared to historical map partly because of swelling of involved fascicles present in patients but not in specimens used by Sunderland.
subsequent improvement, and 3 had abnormal ultrasound examinations at the upper arm. The percentage of true supinator syndrome as a compression neuropathy in patients presenting with finger drop is therefore likely higher than the 16% we observed. Another limitation is that patients with this rare condition were referred from different centers so that clinical examination before the MRI could not be fully standardized.

The study shows that selective proximal RN lesions can account for the clinical appearance of PINS. These lesions can be detected by neuroimaging for accurate lesion localization to complement the basis for correct clinical diagnosis and therapy.

**AUTHOR CONTRIBUTIONS**

Philipp Bäumer: study design, acquisition of data, analysis and interpretation of data, writing of manuscript. Henrich Kele: study design, acquisition of data, analysis and interpretation of data, revising manuscript for intellectual content. Annie Xia: acquisition of data, revising manuscript for intellectual content. Markus Weiler: acquisition of data, analysis and interpretation of data, revising manuscript for intellectual content. Daniel Schwarz: acquisition of data, revising manuscript for intellectual content. Martin Bendszus: study supervision, revising manuscript for intellectual content. Mirko Pham: study design, analysis and interpretation of data, revising manuscript for intellectual content.

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**DISCLOSURE**

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